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(71) Applicant: ALLIED COLLOIDS LIMITED  
P.O. Box 38 Low Moor  
Bradford West Yorkshire, BD12 0JZ(GB)

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(72) Inventor: Farrar, David  
13 Greenfield Lane Idle  
Bradford West Yorkshire(GB)

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(72) Inventor: Hawe, Malcolm  
3 Broombank Birkby  
Huddersfield West Yorkshire(GB)

(74) Representative: Lawrence, Peter Robin  
Broughton et al,  
GILL JENNINGS & EVERY 53-64 Chancery Lane  
London WC2A 1HN(GB)

(54) Polymeric thickeners and their production.

(57) Polymers are made by polymerisation of an ionic monomer such as methacrylic acid or dialkylaminoalkyl (meth)acrylate or -acrylamide and/or a substantially non-ionic monomer, optionally a cross linking agent, and an allyl ether of the formula  $\text{CH}_2=\text{CR}'\text{CH}_2\text{OA}_m\text{B}_n\text{A}_p\text{R}$  where R' is hydrogen or methyl, A is propyleneoxy or butyleneoxy, B is ethyleneoxy, n is zero or an integer, m and p are zero or an integer less than n, and R is a hydrophobic group of at least 8 carbon atoms. The polymers are preferably made by oil in water emulsion polymerisation. They can be used as thickeners particularly in environments containing surfactant and/or electrolyte, including especially emulsion paints, print pastes, alkaline liquors and acidic liquors.

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Allied Colloids Limited

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Polymeric Thickeners and their Production

It is known that aqueous media can be thickened by the presence of high molecular weight synthetic polymers either in solution or in the form of swollen particles.

5 If the polymers are in solution the thickening is probably due to entanglement of polymeric chains. If the polymers are swollen the thickening is probably due to inter-particulate attraction and solvent

10 immobilisation.

It is known that thickening in some instances can be improved by including in the polymer pendant hydrophobic groups, the improvement apparently being due to association between the hydrophobic groups in adjacent

15 molecules and the polymers containing such groups are often referred to as associative thickeners.

GB 1,167,524 describes thickeners that are said to be effective in solutions of surface active agents and that are copolymers of at least one ethylenically

20 unsaturated monomer with a comonomer that is, for instance, an acrylic ester or an allyl ether of a polyalkoxylated alcohol that may be aliphatic or aromatic. This polyalkoxylated alcohol thus provides the pendant hydrophobic groups. Particular allyl ethers

25 of this general type, and copolymers formed from them, are also described in GB 1,273,552.

All the examples in these patents of copolymers using allyl ether show the copolymer to be soluble in water. The comonomer with the ether or ester is said to

30 be acrylic acid, acrylamide, vinyl pyrrolidone or maleic anhydride or a blend of maleic anhydride with a compound which normally copolymerises in alternating sequence. The use of maleic anhydride alone or in such a blend will tend to give a low molecular weight compared to the

values currently available when copolymerising, for instance, acrylic acid.

In most of the examples the comonomer is maleic anhydride, optionally with methyl vinyl ether, but in 5 example 12 of GB 1,167,524 and example 8 of GB 1,273,552 the allyl ether is copolymerised with acrylic acid to form a water soluble polymer. The information in each example suggests that the molecular weight is low.

In all of the examples showing the use of allyl 10 ethers, polymerisation is conducted by precipitation polymerisation. Thus polymerisation is conducted in a liquid that is a solvent for the monomer but not for the polymer, so that polymer is precipitated out of the solution during polymerisation. This technique tends to 15 produce rather low molecular weights.

The intention to form low molecular weight compounds is emphasised by example 13 of GB 1,167,524 in which a copolymer is formed by aqueous solution polymerisation of acrylamide and an acrylic ester with a polyethoxylated 20 nonyl phenyl alcohol since isopropanol is included in the polymerisation mixture. This will ensure that a relatively low molecular weight polymer is produced.

It is well known that allyl monomers polymerise much less readily, and yield copolymers of lower molecular 25 weights, than acrylic or other vinyl monomers. In US 4451628 allyl sulphonate is used to depress molecular weight. Another allyl monomer is diallyl dimethyl ammonium chloride and it is well known that polymers of this generally have a maximum molecular weight of below 30 500,000.

The fact that the use of allyl monomers causes the resultant homo- or co-polymer to be of low molecular weight is discussed frequently in the literature, for instance in "Functional Monomers" Volume 1 edited by 35 Yocum and Nyquist pages 384 to 387. It is stated in

this that allyl monomers polymerise with difficulty to give products of low molecular weight and their presence will retard both the rate and degree of polymerisation of other copolymerisable monomers. It is stated that the 5 polymerisation kinetics of allyl monomers are determined by degradative chain transfer which results in the formation of a stable radical that has low activity for chain propagation. The article describes ways of trying to obtain higher molecular weights but the highest value 10 obtainable is said to have a degree of polymerisation of the order of 1,000 to 10,000. The molecular weight is therefore still low by the standards of acrylic or other vinyl monomers since these can easily be polymerised to molecular weights in the range 10 million to 30 million, 15 provided chain transfer agent is omitted.

Since the techniques and comonomers that were specifically described in GB 1,167,524 were all such as to lead inevitably to relatively low molecular weight polymers it was reasonable to propose the use of allyl 20 ether monomers. They would be expected to give the sort of molecular weights that clearly were intended to be achieved in the processes of that patent and GB 1,273,552. In practice the products of these patents have not proved commercially very successful.

25 In marked contrast to these low molecular weight allyl ether and acrylate copolymers has been the commercial success of associative polymers formed solely from acrylic monomers and by techniques that would normally be expected to give high molecular weights. 30 The literature relating to these polymers generally indicates molecular weights in the range 100,000 (in the presence of chain transfer agent) to several million. Instead of making the polymers in the presence of a relatively large amount of isopropanol or by 35 precipitation polymerisation or using comonomers such as

maleic anhydride blends, all of which will tend to give low molecular weights, the successful associative thickeners are generally made by oil-in-water emulsion polymerisation or by aqueous solution or gel polymerisation, and can have very high molecular weights. If for a particular purpose the highest molecular weights are to be avoided very low amounts of chain transfer agent are incorporated to depress molecular weight but the resultant molecular weights will generally still be well above those made by the processes described in GB 1167524 and 1273552.

In EP 48094 the pendant hydrophobic group is introduced as a polymerisation initiator or chain transfer agent (tending to depress molecular weight) in the polymerisation of acrylamide. In EP 63018 and U.S. 4,423,199 and 4,524,175 the hydrophobic group is introduced as a substituent in acrylamide.

The JP 60-235815A the pendant hydrophobic group is introduced as a vinyl ether.

The great majority of literature on associative thickeners, and all commercial products, introduces the hydrophobic group as an ester of an unsaturated carboxylic acid (generally (meth) acrylic acid) that is copolymerised with one or more monomers that are always vinylic, and are usually (meth) acrylic. Thus in U.S. 3,915,921 and U.S. 4,190,562 the hydrophobic group is introduced as a  $C_{10-30}$  alkyl ester of (meth) acrylic acid. In U.S. 4,138,381, 4,268,641, 4,384,096 and 4,463,151, EP 13836 and EP 109820 and in GB 1,167,524 an ester is formed between an unsaturated acid and a hydrocarbyl ether of a polyalkylene glycol.

When the polymers are linear it is clear that increasing molecular weight generally gives increasing thickening properties (although it may also give flocculation of suspended solids) and so the use of

monomers that make it impossible to obtain high the molecular weights is clearly contra-indicated. In those particular instances where lower molecular weight is desired, e.g., if flocculation of suspended solids is to 5 be avoided, then this is best achieved commercially by using the same monomer blend as will give high molecular weight together with a low amount of a chain transfer agent such as a mercaptan.

Similarly, when the polymers are cross linked 10 experience in other polymerisation techniques for making thickeners generally indicates that the best polymer properties are obtained when the cross linked polymer is formed from monomers that, in the absence of cross linking agent, would give the highest possible molecular 15 weight. Thus cross linked polymers should also be formed from acrylic monomers in the absence of monomers that will significantly reduce molecular weight.

The present state of the art therefore is that when manufacturing cross linked or, especially, linear 20 polymers that are to be used for, for instance, thickening the best properties generally follow from the use of monomers capable of polymerising to very high molecular weights optionally with a chain transfer agent such as a mercaptan, allyl monomers are known to be 25 incapable of giving high molecular weights, and the processes in GB 1,167,524 and 1,273,552 clearly all gave linear polymers of molecular weights much less than those that would now be considered to be necessary for satisfactory properties.

30 Polymers such as those described in EP 13836 are made by oil-in-water emulsion polymerisation and swell or dissolve upon the addition of alkali. They have proved commercially successful but there is still considerable room for improvement in their properties.

For instance one use of the polymers is for thickening aqueous solutions containing an electrolyte. The solutions may therefore have relatively high pH. Also they may be used under conditions of high 5 temperature. Unfortunately high pH and/or high temperature can result in hydrolysis of the ester linkage by which the hydrophobic group is attached.

A problem that is often encountered commercially with thickeners such as in EP 13836 is that they may 10 cause foaming, and print quality (when used in textile printing pastes) may need improvement in some instances.

Despite all the experience indicating that it is essential to use, when making thickeners, only monomers capable of giving high molecular weight polymers, we have 15 now found that a particular class of new polymers formed from monomers including a particular type of allyl monomer have surprisingly valuable thickening properties.

A polymer according to the invention is selected from linear polymers having intrinsic viscosity (as 20 herein defined) above 0.5 dl/g, cross linked polymers and polymers that are insoluble and non swellable in water at pH 7 but that are soluble or swellable in aqueous acid or aqueous alkali and which are formed by polymerisation of  
(a) 0 to 90% of ethylenically unsaturated ionic monomer,  
25 (b) 0 to 90% of ethylenically unsaturated substantially non-ionic monomer,

(c) 0.5 to 100% of ether of the formula  
 $\text{CH}_2=\text{CR}'\text{CH}_2\text{OA}_m\text{B}_n\text{A}_p\text{R}$  where R' is hydrogen or methyl, A is propyleneoxy or butyleneoxy, B is ethyleneoxy, n is zero or an integer, generally 1 to 100, m and p are each zero or an integer less than n and R is a hydrophobic group of at least 8 carbon atoms,  
30 (d) 0 to 5% of cross linking agent, preferably a copolymerisable polyethylenically unsaturated monomer.  
35

Throughout this specification all percentages are by weight unless otherwise specified. All intrinsic viscosities are single point intrinsic viscosity as measured at 0.05% polymer concentration in methanol.

5 The preferred polymers of the invention are made by oil-in-water emulsion polymerisation and are substantially insoluble and non swellable in water at pH 7 but are soluble or swellable in aqueous acid or aqueous alkali and are formed from 5 to 90% of monomer (a), 5 to 10 90% of monomer (b), 0.5 to 90% of monomer (c) and 0 to 5% of monomer (d), and so the invention is now described with particular reference to these.

The oil-in-water emulsion polymerisation is conducted using sufficient of an appropriate emulsifier, 15 as is conventional. The final polymer is insoluble and substantially unswollen in the aqueous phase of the polymerisation mixture but, due to the ionic monomer, is soluble or swellable upon appropriate pH adjustment, generally to below pH 7 when the monomer (a) is free 20 amine and to above pH 7 when the monomer (a) is free acid. The solubility of the monomers in the aqueous phase may be conventional for oil-in-water emulsion polymerisation. Generally the blend of monomers, and often each monomer, is insoluble in the aqueous phase but 25 some water solubility can be tolerated provided the monomers all migrate during the polymerisation into the micelles of emulsifier.

Monomer (a) is preferably a vinyl, generally acrylic, monomer and may be a co-ionic blend of monomers. 30 When monomer (a) is anionic upon addition of alkali, the monomer and its amount must be such that addition of alkali renders the polymer soluble or swellable. The monomer is generally a carboxylic monomer as free acid during the polymerisation. The monomer generally 35 contains 3 to 8 carbon atoms. It may be a

monocarboxylic acid, a dicarboxylic acid or, for instance, a monoalkyl ester of a dicarboxylic acid. The acid may be selected from acrylic, methacrylic, itaconic, crotonic, fumaric, citraconic acryloxypropionic or maleic acids. Preferably at least 50%, and most preferably 100%, of component (a) is provided by methacrylic and/or acrylic acid, with methacrylic being particularly preferred.

When monomer (a) is cationic, the monomer and its amount must be such that the addition of acid or quaternising compound, renders the polymer soluble or swellable. The monomer generally includes a tertiary amine group as a free base during polymerisation and this is then converted to the cationic form as an acid salt or quaternary ammonium salt. Dialkylaminoalkyl (meth) acrylamides may be used. For instance the aminoalkyl group may contain 2-8 carbon atoms, preferably 1,3-propylene, and the other alkyl groups may contain 1 to 4 carbons. The preferred monomers are dialkylaminoalkyl (meth) acrylates. The preferred monomers are dimethylaminoethyl (meth) acrylates and dimethylaminopropyl (meth) acrylamides.

The amount of monomer (a) must be such that the blend of components (a), (b), (c) and (d) can be polymerised by oil-in-water emulsion polymerisation to form an emulsion of the polymeric thickener in which the polymer is insoluble and substantially unswollen and non-thickening but that, after polymerisation, the emulsion can be converted by addition of alkali or acid into a viscous system thickened by the polymer. It is generally necessary for there to be at least 10%, usually at least 20% and preferably at least 30% of the ionic monomer. The amount is generally below 70%, usually below 60%.

Monomer (b) is preferably a vinyl, generally acrylic, monomer and may be a blend of monomers. The monomers are generally water insoluble but a minor proportion of monomer (b) may be a water soluble monomer such as acrylamide. By water insoluble monomer in the context of the present specification we mean monomer that is soluble in water to a degree of up to 5% at room temperature. Suitable monomers are styrene and alkyl- and/or halo-substituted styrenes, (meth) acrylonitrile, vinyl alcanoates (especially the acetate), vinyl and vinylidene halides (especially the chloride), hydroxy alkyl and alkoxy alkyl (meth) acrylates and alkyl (meth) acrylates. Preferred monomers are styrene, 2-hydroxy ethyl acrylate, acrylonitrile, vinyl chloride and vinyl acetate and the alkyl (meth) acrylates. Preferably at least 50% by weight of component (c), and most preferably 100%, is alkyl (meth) acrylate. In all these monomers any alkyl groups may contain 1 to 8 carbon atoms but particularly preferred monomers are C1-4 alkyl (meth) acrylates such as methyl methacrylate, butyl acrylate or, ethyl acrylate.

Monomer (b) is generally present in an amount of at least 15%, usually at least 20% and preferably at least 30%. The amount is generally below 80%, usually below 70% and preferably below 60%.

Normally monomers (a) and (b) are free of hydrophobic groups R and should preferably be conventional low molecular weight monomers.

Monomer (c), which is referred to below as the "allyl ether", preferably includes a polyethoxy chain and so n is generally above 2, often above 5 and frequently above 10 or 15 up to 50 or even up to 100. m and p are each generally zero. Thus a polyoxyethylene chain between the allyl group and the hydrophobe is generally present but it may be interrupted by oxypropylene and/or oxybutylene groups. By appropriate choice of the values of n, m, p and R it is possible to control the solubility

of the monomer and the properties of the final polymer.  
R' is generally hydrogen.

R is a hydrophobic group containing at least 8 carbon atoms. It can be a polyoxyalkylene chain where 5 the alkylene groups wholly or mainly are propylene or butylene or higher but preferably is a hydrocarbyl group.

The hydrocarbyl group generally contains from 8 to 30, preferably 10 to 24 and most preferably 12 to 18 carbon atoms. It may be selected from alkyl, for 10 instance octyl, lauryl or stearyl, aralkyl such as 2-phenyl ethyl (-C<sub>2</sub>H<sub>4</sub>Ph), aryl such as naphthyl, alkaryl such as alkyl phenyl wherein the alkyl group generally contains 6 to 12 carbon atoms, cycloalkyl (including polycyclic alkyl groups), or mixtures of one or more such 15 groups. Preferred hydrocarbyl groups are alkyl and alkaryl groups. Any of these groups may additionally be substituted provided the substituents do not render the pendant group hydrophilic to an extent that the desired improvement in properties due to the hydrophobic group is 20 lost.

The amount of the allyl ether is generally at least 1% and usually at least 2%. It is generally below 70% and usually below 50%. Amounts in the range 5 to 30% are often preferred.

25 The allyl ethers may be made by methods such as those described in GB 1,273,552, for instance by reacting an appropriate surfactant alcohol with sodium or sodium alkoxide, generally in the absence of water but in a solvent such as xylene, to form the sodium derivative and 30 then reacting this with allyl chloride, or by reacting allyl alcohol with the surfactant alcohol with or without catalyst. Preferably however the allyl ethers are made by the method described in our copending application filed even date reference 60/2566.

Monomer (d) is optional and serves as a cross linker. Suitable cross linkers for emulsion polymerised ethylenically unsaturated monomers are well known. They are generally polyethylenically unsaturated monomers 5 materials such as diallyl phthalate, allyl (meth.) acrylate, divinyl benzene, (poly)ethylene glycol dimethacrylate and methylene bis acrylamide. If the cross linker is present its amount is generally in the range 0.0005 to 5% (5 to 50,000 ppm), generally below 1%, 10 most preferably 0.001 to 0.2%.

Particularly preferred copolymers are those formed from 20 to 60% by weight acrylic acid and/or methacrylic acid (preferably methacrylic acid alone), 5 to 60% ethyl acrylate or other suitable alkyl (meth) acrylate and 2 to 15 50% of the allyl ether, optionally with cross linker.

Other preferred polymers are formed from 20 to 60% dialkylaminoalkyl (meth) -acrylate or -acrylamide, 5 to 60% ethyl acrylate or methyl methacrylate or other alkyl (meth) acrylate, with 2 to 50% monomer (c), and 20 optionally (d).

The polymers may have a molecular weight well above 0.5 million and generally above 1 million, eg above 2 million and often above 5 million, when they are prepared in the absence of chain transfer agents. This is far in 25 excess of what would be expected from knowledge of allyl polymerisations in general. However for some purposes the highest molecular weights are undesirable because of the risk of flocculation occurring when the thickener is used in a system containing suspended solids. 30 Accordingly it is sometimes desirable to polymerise in the presence of a chain transfer agent in order to depress molecular weight, eg down to 100000 or 200,000.

The intrinsic viscosity (namely the single point intrinsic viscosity as measured at 0.05% polymer 35 concentration in methanol) of linear polymers is

generally at least 0.5 and when there is no solid phase the IV is preferably at least 1 and preferably at least 2, 3 or even 5, for instance it may be 5-10 or higher.

Cross linked, emulsion polymerised, polymers are 5 polymers made from monomers (a), (b) and (c) under conditions that would, in the absence of cross linker (d), lead to linear polymers having these preferred molecular weight and viscosity properties.

The emulsion polymerised polymers of the invention 10 differ from those of GB 1,167,524 and 1,273,552 in a number of respects. They are made by oil-in-water emulsion polymerisation, this permitting much higher molecular weights. They are insoluble and unswollen at the pH at which they are manufactured, and soluble or 15 swollen at another, whereas the anionic polymers of GB 1,167,524 are water soluble. The molecular weights that are obtained in the invention can be very much higher than anything obtainable in those patents. The reason for our being able to obtain high molecular weights is 20 not clear but may be due to the effect of the group  $-A_m^B_n^A_p^R$  on the polymerisation properties of the allyl monomer. The comonomers preferably are such as to permit high molecular weights and preferably are not such as to reduce molecular weight (e.g., maleic anhydride with 25 methyl vinyl ether, as in those patents).

Other disclosures of polymers containing allyl ethers are in EP 0172723, 0172724 and 0172025, none of which were published at the priority date of this application and all of which, so far as the allyl ether disclosure is concerned, have the same priority date as this application.

The polymers differ from the polymers disclosed in, for instance, EP 13836 by omitting the acrylic ester for introducing the hydrophobic group and using instead the 35 defined allyl ether. It is very surprising that this

substitution can be made without seriously reducing the molecular weight and without damaging the properties of the polymer. It is surprising that the polymers have viscosity and other properties at least as good as those 5 of EP 13836 and in many respects better. By the invention the risk of the hydrophobic group being hydrolysed out of the polymer chain by thermal or pH hydrolysis of the ester linkage is eliminated. It is also possible to avoid the other problems outlined above.

10 The oil-in-water emulsion polymerisation may be conducted in conventional manner, for instance as described in EP 13836. Emulsifier is included to maintain the monomer and the polymer in stable dispersed condition and to provide micelles for the polymerisation.

15 When monomer (a) is anionic, suitable emulsifiers are anionic, such as sodium alkyl phenyl ether sulphate or sodium lauryl sulphate or sodium dodecyl benzene sulphonate, but may be non ionic. When monomer (a) is cationic, the emulsifier is preferably non-ionic.

20 The initiator is preferably a water soluble initiator, most preferably an alkali metal or ammonium persulphate, generally in an amount up to 1% based on the monomers. Preferably polymerisation is started and further monomer is then added. The polymerisation 25 temperature is generally in the range 60 to 100°C.

Although it is generally undesirable, in some instances a chain transfer agent such as an alkyl mercaptan may be added to the monomer in order to depress molecular weight for example to minimise flocculation 30 during thickening.

The amount of monomer, and thus of polymer, in the emulsion is generally from 20 to 60%, most preferably 25 to 50%, based on the weight of emulsion. The emulsion has a pH at which the polymer is insoluble and 35 substantially unswollen in water (7 or below when monomer

(a) is anionic and 7 or above when it is cationic) but upon addition of alkali or acid (for anionic or cationic polymers respectively) the polymer dissolves or swells and will form a very viscous composition. The polymer 5 can be isolated from the water after or, preferably, before the adjustment of pH by, for instance, spray or drum drying to form a powder or the polymer may be converted to a concentrated, substantially anhydrous, dispersion in a non-aqueous liquid as described in EP 10 0172025. Often however it is convenient to use the polymer in the form of the oil-in-water emulsion in which it is initially produced. The pH adjustment may be made to this emulsion before use but preferably the emulsion 15 is added to the phase that is to be thickened while the polymer is still insoluble and unswollen and the polymer is converted to a viscous thickener by reaction with alkali or acid in the aqueous phase. The pH of this aqueous phase may be as high as, for instance, 13 or 14 when the monomer (a) is anionic and as low as 1 or 2 when 20 monomer (a) is cationic.

The emulsion polymerised polymer is generally supplied to the user as an oil-in-water emulsion but if desired may be converted to a water-in-oil dispersion (or a dehydrated product thereof) before use, e.g., as 25 described in EP 172025.

Improved thickening is often achieved in the presence of a surface active agent, generally in an amount of from 0.05 of 1 part by weight surfactant per part by weight polymer. Surfactants that will give this 30 effect may be selected from anionic, non-ionic, amphoteric or cationic provided they are compatible with the polymer and the other components of the aqueous phase. Preferably they are non-ionic or anionic. Preferred surfactants are ethoxylated linear alkyl ethers 35 or ethoxylated alkyl phenyl ethers. Often the

surfactant is the same as or similar to a surfactant alcohol that has been used to form the allyl ether.

The polymers are of particular value for thickening aqueous media containing electrolyte and optionally 5 containing a dispersed phase. For instance the polymers may be used for thickening chemically toxic aqueous compositions. These may be alkaline compositions such as bleaches, caustics and paint removers when monomer (a) is anionic. They may be acidic when monomer (a) is 10 cationic. For instance a cationic polymer may be introduced into an environment and then thickened by adding acid.

The polymers (especially when monomer (a) is anionic) are of particular value for thickening aqueous 15 latex paints, both matt, semi-matt and gloss paints. These may contain conventional pigments, dispersants and binders.

The polymers (especially when monomer (a) is anionic) are also of great value for thickening textile 20 print pastes. Such systems include for example reactive dye or acid dye, or pigment pastes especially where such pastes are prone to gelling. The other components of the print paste may be conventional. In such pastes they not only have the valuable high shear and low shear 25 viscosity properties that are useful in all thickening situations but they also have the advantage of minimising flushing of the print colour. Particularly good results are obtained in combination with surfactant.

Other uses for the polymers are for thickening oils, 30 as water retention aids, eg in cements, and as deicing fluids, for thickening brine (e.g., downhole), for thickening carpet backing latices and for thickening textile sizing solution.

The amount of polymer used is generally in the range 0.05 to 5% weight based on the water content of the aqueous phase that is to be thickened.

The polymers can be used for other purposes, e.g.,  
5 as wallpaper adhesives.

The invention also includes water insoluble, acid or alkali swellable or soluble, polymers made by other techniques, e.g., polymerisation in solution in organic solvent.

10 The invention also includes linear polymers irrespective of how they are made and having IV of at least 0.5. They may be water insoluble and made from the monomer blends and by the methods described above or they may be water soluble. For instance they may be  
15 made from 0.5 to 100% monomer (c) and 0-99.5% water soluble monomers (a) and (b). These may have IV 0.5 to 1 or higher, e.g., as described above.

Monomer (c) may be as described above. The polymers are preferably non ionic or anionic.

20 The preferred soluble polymers have, as monomers (a) and (b), water soluble anionic monomer optionally blended with acrylamide, water soluble cationic monomer optionally blended with acrylamide, or acrylamide. Naturally the monomers should be free of pendant  
25 hydrophobic groups. 0-20%, generally 0-10% and preferably 0%, water insoluble monomer (b) can be included.

One class of preferred soluble anionic copolymers of the invention are formed of 5 to 50%, preferably 20 to  
30 40% acrylic acid or other unsaturated carboxylic or sulphonic acid, generally as a sodium or other salt thereof, 50 to 90%, preferably 60 to 80%, by weight acrylamide and 2 to 30% by weight of the allyl ether.

Another class of preferred soluble anionic copolymers are formed of 50 to 100% (often 80-100%) of

the acid, 0 to 50% (often 0-20%) acrylamide and 2 to 30% of the allyl ether.

Preferred cationic soluble polymers of the invention are formed of 10 to 99% and preferably 20 to 70% dialkyl 5 amino alkyl (meth)-acrylate or -acrylamide quaternary or free acid salt, 0 to 80% preferably 20 to 70% acrylamide, and 1 to 90%, preferably 5 to 50%, of the allyl ether.

The water soluble polymers can be made by conventional methods of making water soluble polymers but 10 modified by the incorporation of the allyl ether, e.g., as in EP 0172723. Thus they may be made by aqueous gel polymerisation or by reverse phase polymerisation. This process may be conducted to a very small dry particle size, such as below 4 µm, for instance as described in 15 European application 0172724 or may be conducted as a bead polymerisation process, using conventional water soluble initiators, stabilisers and, if desired, emulsifiers. Suitable materials are described in that application. Linear water-soluble products may be made 20 as in either of these specifications in the absence of cross-linking monomer.

The soluble polymers may be provided as dispersions in non-aqueous liquid or as dry particles, for instance made by bead polymerisation followed by drying and 25 filtering or made by gel polymerisation followed by drying and comminuting. Depending upon the solubility of the monomers used for their manufacture the polymers will either be truly water soluble or will be water swellable.

These polymers are useful as flocculants, for instance as described in EP 017272. In addition to acting as flocculants for, for instance, sewage or inorganic dispersions they are also of value as filtration aids and paper retention aids, as gangue 35 suppressants as clarification improvers, for pelletising minerals, as dewatering aids or filtration rate improvers as drift controllers in agricultural spray compositions,

as soil stabilisers or dust suppressants, and, especially, as thickeners for aqueous liquids.

They are particularly effective for thickening any of the compositions described above including paints, 5 textile print pastes, or chemically toxic and other aqueous compositions such as bleaches, caustics and paint remover compositions and, especially, for thickening brine, drilling muds and other downhole electrolyte liquors such as for acidising or fracturing, especially 10 when the polymer contains sulphonate groups or cationic groups. Other downhole uses are as viscosifiers for enhanced oil recovery, drilling fluids or shut off fluids, as fluid loss additives, and for polymer flooding. They may be used with surfactant, as 15 described above. The soluble linear polymers may have better suspending properties in respect of large or heavy inorganic particles than the emulsion polymers discussed above.

Another use for the water soluble polymers is as an 20 aqueous adhesive, for instance a wallpaper adhesive that may be a brush-on composition or a prepaste.

The invention also includes cross linked polymers. They may be non swellable in water but swellable in acid or alkali and so may be made by emulsion or organic 25 solution polymerisation as described above. They may be formed of 0.5 to 100% monomer (c) and 0-99.5% monomers (a) and (b), and 0.0001 to 5% cross linking agent, where all the monomers may be as discussed above for emulsion polymerised polymers.

30 Preferably the cross linked polymers are water swellable and the monomers and the polymerisation conditions, are preferably such that in the absence of the cross linking monomer the polymer would have IV at least 1, generally at least 2 and preferably at least 3, 35 often above 5, for instance 10 to 20.

Preferred water swellable polymers are formed from 1 to 100% of the allyl ether, 0 to 99% water soluble monoethylenically unsaturated monomer and 0 to 20% generally 0 to 10% and preferably 0% of water insoluble 5 monoethylenically unsaturated monomer. Suitable water soluble monomers are discussed above.

Preferred swellable polymers have, as monomer (b), water soluble anionic monomer optionally blended with acrylamide, water soluble cationic monomer optionally 10 blended with acrylamide, or acrylamide. Naturally the monomers (b) should be free of pendant hydrophobic groups.

One class of preferred swellable anionic copolymers are formed of 5 to 50%, preferably 20 to 40% acrylic acid 15 or other unsaturated carboxylic or sulphonic acid, generally as a sodium or other salt thereof, 50 to 90%, preferably 60 to 80%, by weight acrylamide and 2 to 30% by weight of the allyl ether, and cross linking agent.

Another class of preferred anionic copolymers are 20 formed of 50 to 100% (often 80-100%) of the acid, 0 to 50% (often 0-20%) acrylamide and 2 to 30% of the allyl ether, and cross linking agent.

Preferred cationic polymers of the invention are formed of 10 to 99% and preferably 20 to 70% dialkyl 25 amino alkyl (meth)-acrylate or -acrylamide quaternary or free acid salt, 0 to 80% preferably 20 to 70% acrylamide, and 1 to 90%, preferably 5 to 50%, of the allyl ether, and cross linking agent.

The water swellable polymers can be made by 30 conventional methods of making water swellable polymers but modified by the incorporation of the allyl ether. Thus they may be made by aqueous gel polymerisation or by reverse phase polymerisation. This process may be conducted to a very small dry particle size, such as

below 4  $\mu\text{m}$ , for instance as described in European application 0172724 or may be conducted as a bead polymerisation process, using conventional water soluble initiators, stabilisers and, if desired, emulsifiers.

5 Suitable materials are described in that application.

The polymers may be provided as dispersions in non-aqueous liquid or as dry particles, for instance made by bead polymerisation followed by drying and filtering or made by gel polymerisation followed by drying and

10 comminuting.

When the polymerisation is conducted to give a small particle size, for instance below 4 microns dry size, and especially when the comonomers are anionic, especially acrylic acid (for instance as ammonium or sodium acrylate) alone or blended with acrylamide the resultant compositions are particularly useful as textile print paste thickeners.

Anionic water swellable polymers having a particle size of, for instance, 50 to 500  $\mu\text{m}$  dry size are 20 particularly valuable as absorbents, for instance in diapers or for dewatering slurries or conditioning soil. Suitable polymers for this purpose are generally formed of 2 to 50% (preferably 3 to 30%) of the allyl ether, 0 to 70% (preferably 30 to 60%) acrylamide and 20 to 98%, 25 preferably 30 to 50%, sodium acrylate and 0.001 to 0.1% cross linking agent. The polymers may be made by gel polymerisation, drying and comminution to the desired size or by reverse phase bead polymerisation.

Alkali swellable, small particle size, polymers from 30 water insoluble monomers are particularly effective for thickening latex paints and chemically toxic and other aqueous compositions such as bleaches, caustics and paint remover compositions and, especially, for thickeneing brine, drilling muds and other downhole electrolyte 35 liquors such as for acidising or fracturing especially

when the polymer contains cationic groups or sulphonate groups. Other downhole uses include viscosifiers for enhanced oil recovery, drilling fluids or shut off fluids, as fluid loss additives, and for polymer flooding.

Other uses for the water swellable polymers is as an aqueous adhesive, for instance a wallpaper adhesive that may be a brush-on composition or a prepaste, as deicing fluids, or as water retention aids.

When the polymer is present as water swellable beads the surface of the particles is preferably less swellable than the inner parts of the particles, preferably as a result of cross linking the surface layer in known manner (e.g., U.S. 3,114,651, 4,043,952 or 4,090,013. This treatment can reduce the stickiness of the particles and can improve the absorption properties.

The following are some examples.

Example 1

An emulsion of monomers in water was prepared by mixing 100g of ethyl acrylate, 80g of methacrylic acid, 20g of allyl ether of 10 mole ethoxylate of stearyl alcohol, 5g of Perlankrol ESD (trade mark), 0.3g ammonium persulphate and 200g water. To a reaction vessel containing 2.5g Perlankrol ESD, 0.1g ammonium persulphate in 255.8g water at 85°C, degassed for 30 minutes with nitrogen there was added 5% of the monomer emulsion over a period of 10 minutes.

After the initial charge had polymerised at 85°C, the remaining monomer emulsion was gradually added over a period of two hours at 85°C. After completion of the monomer feed, the mixture was held at 85°C for 15 minutes and then 10g of 1% ammonium persulphate solution was added. After another 45 minutes, the mixture was cooled and filtered.

The filtrate was approximately 30% solids emulsion copolymer in which the polymer composition is 10% of the allyl ether, 50% ethyl acrylate and 40% methacrylic acid.

The polymer thus prepared was designated Product A.

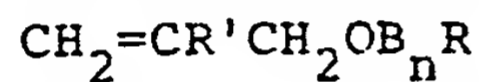
5 Further samples of varying composition were prepared using this procedure. These were designated in turn as Products B-U. Solutions in water were prepared by neutralisation with ammonia to pH 7 or above and the resulting viscosity was measured using a Brookfield RVT  
10 viscometer. This data along with composition variables is given in Tables 1a, 1b, and 1c.

Note In table 1

MAA = methacrylic acid

EA = ethyl acrylate

15 AES = allyl ether surfactant of formula



where R' = H

B =  $(\text{CH}_2-\text{CH}_2-\text{O})$

and n and R are as stated.

20 In table 1a POLYMER H was prepared without using an allyl ether surfactant monomer. This therefore served as a control.

Table 1a

5	PRODUCT	MONOMER RATIOS (% by wt)			SURFACTANT COMPOSITIONS		SOLUTION VISCOSITY (cp) 10rpm
		MAA	EA	AES	n	R	
10	POLYMER A	40	50	10	10	Stearyl	14000
	POLYMER B	40	50	10	2	Stearyl	8400
	POLYMER C	40	50	10	20	Stearyl	7200
	POLYMER D	40	50	10	4	Lauryl	150
	POLYMER E	40	50	10	23	Lauryl	600
	POLYMER F	40	50	10	30	C <sub>22</sub>	140000
15	POLYMER G	40	50	10	25	Octylphenyl	500
	POLYMER H	40	60	-	-	-	60

Note: n - dodecyl mercaptan was included in each of the above recipes, at 0.2% on total monomer.

20

25

30

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Table 1b

5	PRODUCT	MONOMER RATIOS (% by wt)			SURFACTANT COMPOSITIONS		SOLUTION VISCOSITY (cp) 10rpm
		MAA	EA	AES	n	R	
	POLYMER J	40	55.0	5.0	10	Stearyl	1280
	POLYMER K	40	52.5	7.5	10	Stearyl	20800
10	POLYMER L	40	50.0	10.0	10	Stearyl	26200
	POLYMER M	40	47.5	12.5	10	Stearyl	38000
	POLYMER N						
		40	45.0	15.0	10	Stearyl	53600
	POLYMER P	40	42.5	17.5	10	Stearyl	72000
15	POLYMER Q	40	40.0	20.0	10	Stearyl	93200

Note: n - dodecyl mercaptan was included in each of the above recipes at 0.1% on total monomer.

Table 1c

20	PRODUCT	MONOMER RATIOS (% by wt)			SURFACTANT COMPOSITIONS		SOLUTION VISCOSITY (cp) 10rpm
		MAA	EA	AES	n	R	
25	POLYMER L	40	50	10	10	Stearyl	100000
	POLYMER R	30	60	10	10	Stearyl	31000
	POLYMER S	25	65	10	10	Stearyl	15800
	POLYMER T	20	70	10	10	Stearyl	12200
30	POLYMER U	15	75	10	10	Stearyl	260

Note: n - dodecyl mercaptan was included in each of the above recipes at 0.1% on total monomer.

Example 1d

The effectiveness of polymers AA-AF prepared from other alkyl (meth) acrylate/carboxylic acid monomers is shown.

Table 1d

5

Product	Monomer Ratio (% by wt.)								Solution Viscosity (cP) at 10 rpm
	MAA	AA	IA	MA	EA	BA	MMA	AES	
	40	-	-	50	-	-	-	10	
Polymer AA	40	-	-	50	-	-	-	10	76300
Polymer AB	40	-	-	-	-	50	-	10	2400
Polymer AC	35	5	-	-	50	-	-	10	42400
Polymer AD	30	10	-	-	50	-	-	10	11300
Polymer AE	37.5	-	2.5	-	50	-	-	10	48000
Polymer AF	40	-	-	-	25	-	25	10	2940

Note: n-dodecyl mercaptan was included in each of the above recipes at 0.1% on total monomer.

The abbreviations used to denote the monomer are as follows:-

25

MAA = methacrylic acid

AA = acrylic acid

IA = itaconic acid

MA = methyl acrylate

EA = ethyl acrylate

BA = butyl acrylate

MMA = methyl methacrylate

AES = allyl ether surfactant as previously defined with n=10 and R=stearyl.

30

### Example 2

The influence of electrolyte on the thickening efficiency of products of this type was investigated. Three polymers, having compositions as described below were prepared as 2.0% active solutions in deionised water and 0.5% NaCl solution. Solution viscosities were

measured using a Brookfield RVT viscometer over a range of speeds as indicated.

POLYMER V: As POLYMER A but without n-dodecyl mercaptan

5 POLYMER W: As POLYMER v but with 250 ppm diallylphthalate.

POLYMER X: As POLYMER H but with 250 ppm diallylphthalate

Table 2

10

	PRODUCT	2.5 rpm		10 rpm		100 rpm	
		H <sub>2</sub> O	0.5% NaCl	H <sub>2</sub> O	0.5 NaCl	H <sub>2</sub> O	0.5% NaCl
15	POLYMER V 200000	272000		130000	180000	28500	35000
	POLYMER W 284000	240000		180000	85000	25000	22000
	POLYMER X 150000	11200		22000	3700	4200	680

20 Polymer X having no allyl ether surfactant monomer was present to serve as a control.

Polymer V had intrinsic viscosity of 2.5 measured by single point method in methanol at 0.05%.

Example 3

25 The influence of additional surfactant on the thickening efficiency of products of this type was investigated. To 2.0% active solutions of POLYMER A was added increasing amounts of various surfactants. The solution viscosity was remeasured after each addition, and is listed in Table 3 as an index relative to the 30 initial viscosity in the absence of surfactant.

Table 3

	SURFACTANT TYPE	SURFACTANT CONCENTRATION		
		0.1%	0.2%	0.3%
5				
	Lauryl alcohol.4 ethoxylate	188	258	271
	Stearyl alcohol.10 ethoxylate	104	115	135
	Sodium Lauryl sulphate	163	625	798
10	Sodium C <sub>12</sub> -C <sub>15</sub> alcohol ether sulphate	149	214	244

Example 4

To demonstrate the greater stability attained by linking the hydrophobe through an ether linkage rather than an ester linkage, and to eliminate variables due to other monomers in the polymer, a comparison was conducted between the hydrolytic stability of the ester of acrylic acid with polyoxyethylene (23 moles) lauryl ether and of the ether of the same lauryl ether with allyl alcohol (formed by reacting the sodium derivative of the ether with allyl chloride).

A 2.0% solution of each monomer was placed in a thermostated water bath at 45°C. Each was taken to pH 11.0 with 0.0986 M NaOH. The pH was checked every hour and readjusted to pH 11.0 by the addition of further NaOH. The volume of NaOH required was noted and hence the percentage hydrolysis was calculated for each monomer.

Table 4

TIME hours	ACRYLATE MONOMER		ALLYL ETHER MONOMER	
	Vol NaOH(cm <sup>3</sup> )	% Hydrolysis	Vol NaOH(cm <sup>3</sup> )	% Hydrolysis
5	0	1.05	0	1.00
	1	6.75	37.0	0
	2	12.70	69.7	0
10	3	18.10	99.3	0

From this it is apparent that the allyl ether linkage is much more chemically stable than the acrylate linkage.

15 Example 5

In order to demonstrate the usefulness of polymers of this type for thickening aqueous alkaline solution eg caustics and paint removers, solutions of POLYMER A and POLYMER F were prepared 2.0% active in 10% NaOH solution.  
 20 The resulting solution viscosities were measured using a Brookfield RVT viscometer over a range of speeds.

Table 5

PRODUCT	BROOKFIELD VISCOSITY (cp)			
	2.5 rpm	10 rpm	20 rpm	100 rpm
POLYMER A	52000	24500	16000	4600
POLYMER F	54000	16500	9000	2500

30

This clearly demonstrates the high level of thickening efficiency which polymers of this type are able to exhibit in such systems.

Example 6Emulsion Paints

Two matt emulsion paints (MEP1 and 2) were prepared using polymers according to the invention and one (MEP3) 5 using a conventional paint thickener Natrosol 250 HR\*. Each paint was made to 65% pigment volume concentration from the following "Mill Base" and "Let Down" recipes.

<u>Mill Base</u>	<u>Parts by weight</u>
Water	15.00 <sup>1</sup> /19.30 <sup>2</sup>
10 Dispersing agent	0.46
Bactericide	0.05
Hexylene glycol	1.00
Defoamer	0.05
Thickener	1.00 <sup>1</sup> /0.30 <sup>2</sup>
15 0.880 ammonia	0.20
Titanium dioxide	19.75
Calcium carbonate	19.75
Talc	6.59
<u>Let Down</u>	
20 Latex binder	16.18
Coalescing solvent	0.80
Water	18.77 <sup>1</sup> /15.37 <sup>2</sup>
0.880 ammonia	0.40

1. Quantity used with aqueous thickener of the  
25 invention in paints MEP1 and 2.  
2. Quantity used with conventional thickener in paint  
MEP3.

\*Natrosol is a trade mark.

The components of the "Mill Base" were milled under 30 high shear at 2290 rpm to Hegman Gauge 7-8. The components of the "Let Down" were then added at a reduced speed of 890 rpm. The paints were then stored for a period of 7 days at room temperature before characterisation.

Table 6

	Paint Sample	Thickener (0.3% dry Code on total)	Final pH	Brookfield RVT @ 20 rpm (Poise)	Stormer Viscosity (Krebs Units)	ICI Rotothinner Viscosity (Poise)
5	MEP-1	POLYMER A	9.0	120	108	9.5
10	MEP-2	POLYMER W	9.0	252	121	10.0
15	MEP-3	NATROSOL 250HR	9.1	120	105	7.6

Each of the above points was evaluated for its spatter properties using the following test procedure.

15 A board, approximately 30 x 42 cm was mounted vertically above a work bench the bottom edge being positioned 20 cm above the surface of the bench. A piece of black card measuring 24 x 31.5 cm was then placed on the surface of the bench top directly below the board in 20 order to catch any droplets or "paint mist" spattering from the roller.

25 Each paint was poured in turn into a roller tray and applied to the vertical board using a lambswool roller. The standard conditions adopted for this test involved 10 upstrokes and downstrokes of the roller on the board. The black card was then removed for inspection. The degree of spatter was assessed visually and rated on a scale of 1-5 with 1 being excellent; ie no spatter and 5 being very poor; ie severe spatter. The results recorded 30 were as follows:

<u>Paint Sample Code</u>	<u>Spatter</u>
MEP-1	2 - Very Good
MEP-2	1 - Excellent
MEP-3	4 Poor

35 Example 7

Carpet backing formulations

The use of products of this type as thickeners for carpet backing formulations has been demonstrated according to the following information.

	<u>Formulation</u>	A	B
5	Total Solids (nominal)	76%	40%
	Filler: Binder Ratio (dry:dry)	8:1	1:2
	Latex Binder (50% active)	15.6	49.5
	Calcium carbonate filler	67.5	13.0
10	Surfactant	0.1	0.1
	Dispersant	1.0	1.0
	Thickener + Water	to 100	to 100

The amount of thickener was selected such that Formulation A had viscosity 9000cp and B had viscosity 15 6000cp, Brookfield RVT viscometer, spindle 5 at 10 rpm.

In this instance Polymer E fully neutralised as  $\text{Na}^+$  salt was compared as thickeners with VISCALEX AH 10 (carboxylated acrylic gel polymer sold by Allied Colloids Limited as a thickener for carpet backing compounds).  
20 The following results were obtained.

Table 7

BROOKFIELD VISCOSITY (CP)				
5	FORMULATION A	FORMULATION B		
THICKENER TYPE	Viscalex AH10	Polymer E	Viscalex AH10	Polymer E
ADDITION LEVEL	0.16	0.035	0.90	0.15
10 (‡ Dry on Total)				
Initial Viscosity	9100	8600	5800	6000
Day 1 Viscosity	9000	8600	5600	6200
15 Day 2 Viscosity	9000	8600	5800	6200

The improved level of efficiency of the polymers of the invention is clearly demonstrated.

#### Printing Pastes

20 In the examples which follow, all print paste viscosities were measured with a Brookfield RVT viscometer at speed 10 rpm, spindle 6 at 20°C. Printing was carried out using a variety of screens, each being constructed of 156 mesh polyester filament.

#### Example 8

A printing clear was prepared according to the recipe given in Table 8.

Table 8

COMPONENT	% by wt
5 Ammonia (0.880)	0.5
Emulsifier*	2.0
Odourless kerosene	40.0
Binder latex (40% solids)	12.0
Polymer W	1.7
10 Water	43.8

\* e.g., ethoxylated nonyl phenol (14 moles ethylene oxide)

A viscosity of 20,000 cp was obtained after stirring 15 for 10 minutes. The clear was then divided and pigment printing pastes prepared by mixing 9.6 parts of this stock with each of the following pigments:

- (i) Imperom Blue KRR - 4 parts
- (ii) Helizarin Bordeaux R - 4 parts

20 These pastes were then re-thickened by stirring in additional quantities of Polymer W until a viscosity of 20,000 cp was again obtained. The amounts of Polymer W required were found to be 0.19 parts and 0.23 parts for the blue and bordeaux pigments respectively.

25 Both pastes were printed on to a plain woven 50/50 cotton fabric. The prints were then dried and cured for 4.5 minutes of 150°C. In each case excellent colour yield, brightness and fine line definition were observed.

#### Example 9

30 A series of printing clears was prepared in which the concentration of Polymer A was progressively increased. The recipes are given in Table 9a below, where all figures refer to parts by weight:

Table 9a

Component	Paste	A	B	C	D
5					
Ammonia (0.880)		0.5	0.5	0.5	0.5
Binder latex (40% solids)		12.0	12.0	12.0	12.0
Water		85.88	85.65	85.56	85.21
10 Thickener <sup>+</sup>		1.62	1.60	1.44	1.29
Polymer A		-	0.25	0.50	1.00

15           <sup>+</sup> The thickener used was a 50% active dispersion of  
a cross-linked polyacrylic acid such as described in  
US 4554018. In each case the concentration employed was  
chosen so as to give a viscosity of 20,000 cp.

20           Printing pastes were then prepared from each clear  
by mixing 96 parts with 4 parts of the following black  
pigments and stirring by efficient mechanical means for  
5 minutes.

- (i) Imperon Black KGF
- (ii) Acramin Black FBRK

The viscosity of each paste was then remeasured at  
20°C. The results were as follows:

25

30

35

Table 9b

PASTE	VISCOSITY (cp)	
	Black KGF	Black FBRK
5		
A	50,000	66,000
B	29,400	30,600
C	23,000	25,200
10 D	19,600	20,200

The two pastes A, prepared without the addition of Polymer A, were highly viscous gels and were considered unsuitable for practical use. As the concentration of 15 Polymer A was increased, the increase in viscosity, compared with the initial clears, was greatly reduced. All pastes B, C and D were of the correct rheology for commercial printing.

Example 10  
20 Printing pastes were prepared using Polymer A and a cross-linked polyacrylic acid thickener in an analogous manner to that outlined above. The recipes were as follows, all figures referring to parts by weight.

25

30

35

Table 10a

Paste Component	A	B	C	D	E	F	G	H
Ammonia (0.880)	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
Binder latex (40%)	12.0	12.0	12.0	12.0	12.0	12.0	12.0	12.0
Water					to 100%			
Thickener	1.62	1.61	1.44	1.29	1.65	1.84	1.82	1.79
Polymer A	-	0.25	0.5	1.0	-	0.25	0.5	1.0
Imperon Blue KRR	4.0	4.0	4.0	4.0	-	-	-	-
Helizarin								
Bordeaux R	-	-	-	-	4.0	4.0	4.0	4.0

Each paste was printed on to a substrate which was known to cause severe flushing or haloing when printed with many existing formulations. The fabric chosen was a plain woven 50/50 polyester/cotton which had been preresinated using a DHDHEU resin and also contained a magnesium chloride catalyst.

The severity of flushing was critically assessed by using a printing screen which consisted of a series of five lines and criss-cross patterns. Immediately after printing, the fabric was allowed to dry for 2 minutes under atmospheric conditions before being dried at 150°C for 2 minutes (drying quickly causes the movement of print paste to be halted).

The extent to which flushing occurred was then assessed on an arbitrary 1-10 scale, where 1 indicates a perfect print without any sign of flushing and 10 indicates severe flushing. The results are given below:

Table 10b

	PRINT PASTE	BLUE	PASTE	BORDEAUX
5	A	8	E	8
	B	5	F	3
	C	3	G	3
	D	2	H	2

10 The inclusion of Polymer A into the printing paste is clearly seen to bring about a dramatic improvement in print quality, with a 1% addition virtually eliminating flushing. It is considered that a rating of 3 or below would represent a commercially acceptable print.

15 Example 11

A latex was formed by oil in water emulsion polymerisation using ammonium persulphate of 40 parts by methacrylic acid, 50 parts by weight ethyl acrylate and 10 parts by weight of a surfactant ether formed from 20 allyl chloride and the reaction product of polyoxyethylene (10 moles) stearyl ether with sodium methoxide.

25 200 grams of the latex was homogenised into an oil phase comprising 7.0 grams of Span 80, 23.3 grams of a 30% active solution of a 2:1 molar copolymer of ceto-stearyl methacrylate:hydroxyethylmethacrylate in SBP11, 39.2 grams of Pale Oil 60 and 96.0 grams of SBP11. The resulting inverse emulsion was then dehydrated by distilling off water and SBP11 under reduced pressure to 30 a final pressure of 10 mm.Hg and a temperature of 95°C.

The resulting anhydrous polymer-in-oil dispersion was activated by mixing in 5.6 grams of Ethylan D254 to produce a self-emulsifiable liquid polymer-in-oil dispersion having 50% active solids. Span and Ethylan 35 are trade marks.

Upon addition of this dry dispersion to deionised water or 0.5% NaCl solution to form a 2% solution, the Brookfield viscosity at spindle 6 is 352,000 or 480,000 cps respectively at 2.5 rpm and 28,000 or 35,000 cps at 5 100 rpm.

Example 12

The process of Example 1, for Polymer A, was repeated except that the free acid monomer was replaced by a free base monomer as indicated in Table 12a. The 10 Perlankrol surfactant was replaced by Ethylan HA (non-ionic).

Polymers having the compositions indicated in the table below were prepared at 1% active in water by neutralisation with HCl, and the resulting viscosity was 15 measured using a Brookfiled RVT viscometer.

Table 12a

Product	Monomer Ratio (% by wt.)		Composition	Surfactant		Solution Viscosity (cP) @ 10 rpm
	DMAEMA	NMA		AES	n	
Polymer AH	45	35	20	10	stearyl	29000
Polymer AJ	50	30	20	10	stearyl	68000
Polymer AK	55	25	20	10	stearyl	136000
Polymer AL	60	20	20	10	stearyl	80000
Polymer AM	40	35	25	10	stearyl	74000
Polymer AN	40	30	30	10	stearyl	27000
Polymer AP	40	25	35	10	stearyl	5200
Polymer AQ	40	20	40	10	stearyl	1700

DMAEMA = dimethylaminoethylmethacrylate

MMA = methylmethacrylate

AES = allyl ether surfactant

In order to demonstrate the effectiveness of polymers of this type in thickening highly acidic media Polymer AH was used to thicken 15% solutions of different acids at 5% active polymer. The results obtained are 5 given in Table 12B.

Table 12b

10	Product	Acid Type (solution conc <sup>3</sup> w/w)	Solution Viscosity (cP) @ 10 rpm
	Polymer AH	15% hydrochloric acid.	2,200
	Polymer AH	15% phosphoric acid	6,400
	Polymer AH	15% acetic acid	9,600
15			

This polymer is useful for thickening aqueous acidic liquors, such as battery liquids.

Example 13

20 181.8 parts of a 79.2% solution of acrylic acid in water, 0.4 parts of Tetralon B, 5.8 parts of the allyl ether of a 10 mole ethoxylate of stearyl alcohol, 118 parts of water, 0.0424 parts of AZDN and 116 parts of a 29.9% solution of ammonia in water were mixed to form an 25 aqueous solution. A non-aqueous liquid phase was formed from 7.4 parts of Span 80, 42.4 parts of a 30% solution in SBP11 of an inverse dispersion stabiliser (copolymer of 2 moles cetostearyl methacrylate with 1 mole of methacrylic acid), 127.3 parts of Pale Oil 60 and 145.7 30 parts of SBP11.

The aqueous phase was homogenised into the oil phase, deoxygenated and polymerised using 0.042 parts of sodium metabisulphite dissolved in 2.058 parts of water and tertiary butyl hydroperoxide added continuously as a

0.5% solution in water at a rate of 0.14 parts per minute.

The resulting inverse dispersion of hydrated polymer was distilled to yield a dehydrated concentrated polymer dispersion to which was added 2 parts of a 5 mole ethoxylate of nonyl phenol and 1 part of a 4 mole ethoxylate of a broad cut lauryl alcohol per 100 parts of concentrated dehydrated dispersion.

This formed a dispersion of 50% active copolymer which dispersed with agitation in water to yield a highly viscous polymer solution with the characteristic 'soap gel' rheology of associated water soluble polymers. This process was repeated except that the aqueous phase contained in addition 0.063 parts of methylene bis acrylamide as bi-functional crosslinking comonomer. The resultant dehydrated polymer particles swelled in water to form a highly viscous but non-viscoelastic paste useful as a vehicle for printing textiles and other articles particularly on cloth containing residual electrolyte where pastes thickened with conventional polyammonium acrylate microgel latices give holoing, bleeding or flushing of print colour.

Polymers produced by this method but omitting the cross-linking monomer may be made and may be as exemplified in EP-A-0172724.

#### Example 14

A range of polymers were made by gel polymerisation using differing amounts of acrylamide, sodium acrylate, allyl ether and cross linking agent, and by using different allyl ethers. The product of the polymerisation was then dried and comminuted to give particles having sizes in the range 200 to 500 microns.

0.5 g of each polymer was added to 400 cc of a swelling solution that was deionised water or aqueous sodium chloride of various concentrations. The samples

were allowed to equilibrate for 30 minutes and the swollen gel particles were then separated from the medium by filtration through a nylon filter mesh and weighed to give an indication of absorbency. The values are  
 5 expressed in percentage based on the amount of deionised water that was absorbed by each polymer. The monomer feed and the results are shown in the following table. In this table R is the hydrophobic group and n is the number of ethylene oxide groups between it and the allyl  
 10 ether linkage.

Polymer	1	2	3	4	5	6	7	8
Acrylamide	60	50	45	45	50	50	50	50
Sodium Acrylate	40	40	40	40	40	40	40	40
Methylene bis acrylamide	0.03	0.06	0.06	0.06	0.03	0.06	0.03	0.06
Allyl ether	0	10	15	15	10	10	10	10
R	- C <sub>18</sub> H <sub>37</sub>	C <sub>18</sub> H <sub>37</sub>	C <sub>18</sub> H <sub>37</sub>	C <sub>12</sub> H <sub>25</sub>	C <sub>12</sub> H <sub>25</sub>	C <sub>18</sub> H <sub>37</sub>	C <sub>18</sub> H <sub>37</sub>	
n	-	10	10	20	23	23	20	20
Deionised water	100	100	100	100	100	100	100	100
0.1% NaCl	21.7	32.4	25.5	33.8	27.3	21.8	25.1	34.0
0.25% NaCl	12.5	22.4	16.7	22.9	18.8	14.9	16.6	23.8
0.5% NaCl	10.3	16.5	11.6	14.9	15.1	11.6	11.9	17.4
1% NaCl	7.9	8.8	10.0	11.2	11.6	7.9	9.8	13.6

30 Comparison of polymers 3 and 4 shows the advantage that follows from increasing the length of the ethoxy chain. Very poor results are obtained when there is no ethoxy chain, as in U.S. 4,190,562.

Linear water-soluble polymers may be made as in this  
 35 example but omitting the cross-linking monomer. Thus they may be as exemplified in EP-A-0172723.

CLAIMS

1. A polymer selected from polymers that are substantially non swelling and insoluble in water at pH 7 but soluble or swellable in aqueous acid or alkali, cross linked polymers, and linear polymers having intrinsic viscosity at least 0.5 (measured as single point at 0.05% concentration in methanol) and formed by polymerisation of
  - (a) 0 to 90% by weight of ethylenically unsaturated ionic monomer
  - (b) 0 to 90% by weight of ethylenically unsaturated substantially non-ionic monomer
  - (c) 0.5 to 100% by weight of an ethylenically unsaturated monomer that carries a pendant group - $A_m B_n A_p R$  where A is propylenoxy or butylenoxy, B is ethylenoxy, n is zero or a positive integer, m is zero or an integer less than n and R is a hydrophobic group of at least 8 carbon atoms
  - (d) 0 to 5% by weight of cross linking agent, characterised in that monomer (c) is an allyl ether of the formula  $CH_2 = CR'CH_2OA_m B_n A_p R$  where R' is hydrogen or methyl.
2. A polymer according to claim 1 in which monomer (a) comprises a vinyl carboxylic acid monomer.
3. A polymer according to claim 2 in which monomer (a) is selected from acrylic acid and methacrylic acid and mixtures thereof.
4. A polymer according to claim 1 in which monomer (a) comprises a vinyl tertiary amine monomer.
5. A polymer according to claim 4 in which monomer (a) comprises a dialkylaminoalkyl (meth) acrylate monomer or dialkylaminoalkyl (meth) acrylamide.
6. A polymer according to any preceding claim in which monomer (b) is selected from styrene, alkyl-substituted

styrenes, halo-substituted styrenes, (meth) acrylonitrile, vinyl alkanoates, vinyl halides, vinylidene halides, hydroxy alkyl (meth) acrylates, alkoxy alkyl (meth) acrylates and alkyl (meth) acrylates.

5 7. A polymer according to claim 6 in which monomer (b) is selected from Cl-4 alkyl (meth) acrylates.

8. A polymer according to any preceding claim in which, in monomer (c), R' is hydrogen, m and p are each zero and n is from 5 to 100.

10 9. A polymer according to any preceding claim in which R is selected from hydrocarbyl groups and polyoxyalkylene chains wherein the alkylene groups are wholly or mainly propylene or butylene.

10. A polymer according to any preceding claim in which, 15 in monomer (c), R is a hydrocarbyl group of 8 to 30 carbon atoms selected from alkyl, aralkyl, aryl, alkaryl and cycloalkyl.

11. A composition according to any preceding claim in which, in monomer (c), R contains 10 to 24 carbon atoms 20 and is selected from alkyl and alkaryl.

12. A composition according to claim 1 made by oil in water polymerisation of 5-90% monomer (a), 5-90% monomer (b), 0.5-90% monomer (c) and 0-5% cross linking agent.

13. A polymer according to claim 12 formed from 20 to 25 60% by weight acrylic acid and/or methacrylic acid, 5 to 60% by weight ethyl acrylate and 2 to 50% by weight of the allyl ether, and 0 to 5% cross linking agent.

14. A polymer according to claim 1 in which the polymer is linear, is formed in the absence of the cross linking 30 agent, and has a molecular weight above 0.5 million.

15. A polymer according to claim 14 that is water soluble and formed from 5-98% water soluble ionic or non ionic monomers and 2 to 95% monomer (c).

16. A composition according to claim 1 in which the polymer was formed in the presence of a cross linking agent.
17. A polymer according to claim 8 that is water swellable and formed from 5-95% water soluble ionic or non ionic monomers and 2-95% monomer (c).
18. A polymer according to claim 15 or claim 16 formed from acrylic acid or dialkylaminoalkyl (meth)-acrylate or -acrylamide, and 0-80% acrylamide, and 2 to 50% of monomer (c).
19. A polymer according to claim 15 or claim 16 that is anionic.
20. A composition comprising a polymer according to claim 12 which is in the form of an aqueous emulsion at a pH such that the polymer is insoluble and substantially unswollen and non thickening but can be converted to a dissolved or swollen and thickening state by the addition of acid or alkali.
21. A composition comprising a polymer according to claim 12 and that is at a pH such that the polymer is dissolved or swollen and the composition is thickened by the polymer.
22. A composition according to claim 21 selected from print pastes and emulsion paints.
23. A composition according to claim 20 or claim 21 and which also contains surfactant in an amount of 0.05 to 1 part per part by weight polymer.



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EUROPEAN SEARCH REPORT

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DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document with indication where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int Cl *)
A	US-A-4 384 096 (L.F. SONNABEND) * Claim 1 *	1	C 08 F 246/00 C 08 F 220/28 C 08 F 216/14 C 09 D 7/00
A	FR-A-2 007 379 (TOYO RAYON) * Claim 1 *	1	
A	EP-A-0 003 235 (BASF) * Claim 1 *	1	
A	EP-A-0 056 627 (NIPPON SHOKUBAI KAGAKU KOGYO CO. LTD.) -----	1	
TECHNICAL FIELDS SEARCHED (Int Cl *)			
C 08 F			
The present search report has been drawn up for all claims			
Place of search	Date of completion of the search	Examiner	
THE HAGUE	28-11-1986	CAUWENBERG C.L.M.	
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